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1     **Cellulose propionate/poly(*N*-vinyl pyrrolidone-*co*-vinyl acetate) blends: dependence of**  
2     **the miscibility on propionyl DS and copolymer composition**

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**Abstract:** Blend miscibility of cellulose propionate (CP) with synthetic copolymers comprising *N*-vinyl pyrrolidone (VP) and vinyl acetate (VAc) units was examined, and a data map was constructed as a function of the degree of substitution (DS) of CP and the VP fraction in the copolymer component. Results of DSC and FT-IR measurements indicated that the pairing of CP/P(VP-*co*-VAc) formed a miscible or immiscible blend system according to the balance in effectiveness of the following factors: 1) hydrogen bonding between residual hydroxyls of CP and VP carbonyls of P(VP-*co*-VAc); 2) steric hindrance of propionyl side-groups to the interaction specified in 1); 3) intramolecular repulsion between the two units constituting the vinyl copolymer; and, additionally, 4) structural affinity between two segmental moieties involving the propionyl group and VAc unit, respectively. The factor 3 inducing intercomponent attraction is responsible for the appearance of a so-called "miscibility window" in the miscibility map, and the factor 4 substantially expands the miscible region whole, wider relative to those in the maps for the corresponding blend series based on cellulose acetate and butyrate. In further refined estimation by DMA and  $T_{1\rho}^H$  quantification in solid-state  $^{13}\text{C}$  NMR, it was found that the miscible blends of hydrogen-bonding type (using CPs of  $\text{DS} < 2.7$ ) were completely homogeneous on a scale within a few nanometers, whereas the polymer pairs situated in the window region (using CPs of  $\text{DS} > 2.7$ ) formed blends exhibiting a somewhat larger size of heterogeneity (ca. 5–20 nm).

**Keywords:** Blends; Cellulose propionate; Poly(*N*-vinyl pyrrolidone-*co*-vinyl acetate); Miscibility; Scale of homogeneity

## Introduction

As is well known, polymer blending is useful to improve the original physical properties of one or both of the components, and also to obtain new polymeric materials exhibiting wide-ranging properties and/or synergistic functions unattainable in single-component materials (Ultracki 1990). This should also be applicable to the blending of cellulosics as one component (Nishio 1994). Especially cellulose esters (CEs) are versatile cellulosic derivatives and essential for further applications in various fields including molded plastics, fibers, optical films, membranes, coatings, etc., and, therefore, a number of fundamental and practical blend studies of CEs have been carried out (Edgar et al. 2001; Nishio 2006).

In previous papers (Miyashita et al. 2002; Ohno et al. 2005; Ohno and Nishio 2006), the authors' group has investigated the miscibility and intermolecular interactions for blends of industrially crucial CEs, cellulose acetate (CA) and butyrate (CB), with synthetic homo- and copolymers comprising *N*-vinyl pyrrolidone (VP) and/or vinyl acetate (VAc) units, i.e., poly(*N*-vinyl pyrrolidone) (PVP), poly(vinyl acetate) (PVAc), poly(*N*-vinyl pyrrolidone-*co*-vinyl acetate) (P(VP-*co*-VAc)). Through thermal analysis by differential scanning calorimetry (DSC), it was shown that the miscibility behaviour of the CA or CB/vinyl polymer pairs (generically described as CE/P(VP-*co*-VAc)) was seriously affected by the degree of substitution (DS) and the ester side-chain length of the CE component, as well as by the VP fraction in the copolymer component. Two maps given in Figure 1 survey the estimation result.

### <<Figure 1 (a) & (b)>>

In the CA/P(VP-*co*-VAc) system (Fig. 1a), Fourier transform infrared (FT-IR) and solid-state <sup>13</sup>C CP/MAS NMR spectroscopy revealed that the blend miscibility was mainly governed by the hydrogen-bonding interactions between the residual hydroxyls of CA and the carbonyls of VP units in P(VP-*co*-VAc) and the miscible blends of this interaction type were



homogeneous in a few nanometers scale (Miyashita et al. 2002; Ohno et al. 2005). In the CB/P(VP-*co*-VAc) system (Fig. 1b), the hydrogen-bonding interaction was suppressed in frequency by steric hindrance of the bulky butyryl substituent, resulting in lowering of the critical DS required for attainment of the miscibility of CE with PVP and VP-rich copolymers (VP content > 65 mol%), as the critical values of 2.5 for CB and 2.8 for CA are designated in Figure 1. Furthermore, unlike the situation for the CA blends, highly substituted CBs of DS = 2.5–2.95 made a miscible pair with P(VP-*co*-VAc) copolymers containing ca. 30–65 mol% VP residues (Ohno and Nishio 2006). This unique copolymer composition range, generally termed a ‘miscibility window’, emerges as a result of indirect polymer-copolymer attraction driven by strong repulsion between the VP and VAc constituents of the random copolymer. More concretely, since these two monomer species having mutually repellent characters were randomly combined in P(VP-*co*-VAc) by covalent bonding, the copolymers tended to form a miscible monophasic system with CB (DS > 2.5) so as to reduce the strong repulsion between the comonomers (Ohno and Nishio 2007). The absence of such a clear miscibility window in the map for the CA/P(VP-*co*-VAc) system may be interpreted as due to a strong self-association ability of highly substituted CAs of DS > ~2.8; the CAs rather crystallize in a cellulose triacetate II form.

As an extension of the above studies, our attention was then directed to a similar miscibility map for cellulose propionate (CP)/P(VP-*co*-VAc) blends; the side-chain length of the CE component is just intermediate between the acetyl and butyryl substituents. Great interests are how far the miscible region spreads on the map constructed as a function of the DS and copolymer composition, and whether that kind of miscibility window emerges or not. Thereby, we will be able to make clearer the effects of the ester side-group and residual hydroxyls of CE on the blend miscibility and intermolecular interactions with the vinyl polymers concerned. In addition to conventional characterizations by DSC analysis and IR and NMR spectra, the homogeneity of miscible blends is evaluated in refinements of the

mixing scale by complementary use of dynamic mechanical analysis (DMA) and proton spin-lattice relaxation time ( $T_{1\rho}^H$ ) measurements in solid-state  $^{13}\text{C}$  NMR spectroscopy.

## Experimental

### Materials

Cellulose propionate (CP) samples were synthesized from cotton cellulose with a viscosity average molecular weight of 252,000 via a homogeneous reaction with acid chloride/base catalyst, in a procedure similar to that used in previous studies (Kusumi et al. 2008; Nishio et al. 1997). Table 1 summarizes the characterization data including DS, molecular weight, and glass transition temperature ( $T_g$ ) for all the CP samples used in this study. The vinyl polymers employed as a mixing partner for the CPs were poly(*N*-vinyl pyrrolidone) (PVP), poly(vinyl acetate) (PVAc), and poly(*N*-vinyl pyrrolidone-*co*-vinyl acetate) (P(VP-*co*-VAc)), basically the same as those in the preceding papers (Miyashita et al. 2002; Ohno and Nishio 2006). Data of characterization for all the vinyl polymers are also listed in Table 1. As shown in the table, any of the P(VP-*co*-VAc) samples exhibited a single  $T_g$ , and the  $T_g$  versus copolymer composition relation was in good obedience to the Fox equation (Fox and Flory 1954). Thus the copolymers were all regarded as essentially random copolymer. Hereafter, a CP sample with  $\text{DS} = x$  is encoded as  $\text{CP}_x$ , and a code P(VP<sub>*y*</sub>-*co*-VAc<sub>*z*</sub>) denotes P(VP-*co*-VAc) copolymer of VP:VAc = *y*:*z* (in molar ratio).

<<Table 1>>

### Preparation of blend samples

CP/vinyl polymer blends were prepared in film form from mixed polymer solutions by

solvent evaporation, in the same manner as that adopted in the preceding works (Miyashita et al. 2002; Ohno and Nishio 2006). *N,N*-Dimethylformamide was selected as a common solvent and the film casting was carried out at 50 °C under reduced pressure (< 10 mmHg). The as-cast samples thus obtained were further dried at 50 °C *in vacuo* for 3 days.

For DMA measurements, the solution-cast samples were thermally molded into a flattened film ca. 0.1 mm thick by using a Toyo-Seiki hot-pressing apparatus. The hot-press molding was conducted at 230 °C with an applied pressure of 15 MPa for 30 s.

## Measurements

DSC thermal analysis was carried out with a Seiko DSC 6200/EXSTAR 6000 apparatus. The temperature readings were calibrated with an indium standard. The calorimetry measurements were conducted on ca. 5-mg samples packed in an aluminum pan under a nitrogen atmosphere. Each sample was first heated from ambient temperature (~25 °C) to 230 °C at a scanning rate of 20 °C/min, and then immediately quenched to –50 °C at a rate of 80 °C/min. Following this, the second heating scan was run from –50 °C to 230 °C at a rate of 20 °C/min to record stable thermograms. Thermograms presented in this paper were all obtained in the second heating scan and the  $T_g$  was taken as a temperature at the midpoint of a baseline shift in heat flow characterizing the glass transition.

FT-IR spectra were measured on thinner film samples (<20 μm thick) by using a Shimadzu IRPrestige-21 spectrometer. All the spectra were recorded at 20 °C in a transmission method over a wavenumber range 400–4000 cm<sup>–1</sup> with a resolution of 2 cm<sup>–1</sup> via accumulation of 64 scans.

DMA was conducted by using a Seiko DMS6100/EXSTAR6000 apparatus. Strips of rectangular shape (20 × 5 mm<sup>2</sup>) cut from the molded films were used for measurements of the temperature dependence of the dynamic storage modulus ( $E'$ ) and loss modulus ( $E''$ ). The

measuring conditions were as follows: temperature range,  $-150$ – $300$  °C; scanning rate,  $2$  °C/min; oscillatory frequency,  $10$  Hz.

High-resolution solid-state NMR experiments were performed at  $20$  °C in a Varian NMR system  $400$  MHz operated at a  $^{13}\text{C}$  frequency of  $100.6$  MHz. The magic-angle spinning rate was  $15.0$  kHz.  $^{13}\text{C}$  CP/MAS spectra were measured with a contact time of  $2$  ms, and a  $90^\circ$  pulse width of  $2.9$   $\mu\text{s}$  was employed. In the measurements of  $T_{1\rho}^{\text{H}}$ , a contact time of  $0.2$  ms was used, and a proton spin-locking time  $\tau$  ranged from  $0.5$  to  $30$  ms.  $2048$  scans were done to obtain the  $^{13}\text{C}$  CP/MAS spectra, while  $4096$  scans were accumulated for the relaxation time measurements. Chemical shifts of  $^{13}\text{C}$  spectra represented in ppm were referred to tetramethylsilane by using the methine carbon resonance ( $29.47$  ppm) of adamantane crystals as an external reference standard. In order to minimize any possible effect due to the thermal history and/or residual solvents, each sample was heat-treated at  $250$  °C *in vacuo* for  $5$  min just before the measurement.

## Results and discussion

### Estimation of miscibility and intermolecular interaction

The miscibility state in the present CP/vinyl polymer system was estimated basically by  $T_g$  determination in DSC; generally, if any blend sample of a given polymer/polymer pair exhibits a single glass transition between the  $T_g$ s of the two component polymers and a composition-dependent shift of the blend  $T_g$  is clearly observed, then the pair can be regarded as a miscible one on the  $T_g$ -detection scale that is usually assumed to be less than a couple of tens of nanometers (Kaplan 1976; Nishio 1994; Ultracki 1990). To examine the presence of intermolecular interactions, different blend compositions of selected CP/vinyl polymer pairs were subjected to FT-IR and CP/MAS NMR spectra measurements.

163

## 164 *CP/PVP blends*

165 When CPs of  $DS = 1.71\text{--}2.62$  were used as a counter component to PVP, the solution-cast  
166 blend films prepared at 10/90–90/10 (wt/wt) compositions were all transparent in the visual  
167 inspection. By contrast, CP/PVP blends of propionyl  $DS = 2.72\text{--}2.93$  formed a  
168 comparatively cloudy film at intermediate compositions of 40–70 wt% CP content.

169 Figure 2a displays DSC thermograms obtained for  $CP_{2.72}/PVP$  blends. From reading of  
170 the midpoint of the respective discontinuities in heat flow,  $T_g$  of  $CP_{2.72}$  and that of PVP were  
171 evaluated as  $134\text{ }^{\circ}\text{C}$  and  $177\text{ }^{\circ}\text{C}$ , respectively. For the blend samples of 20/80–80/20  
172 compositions, two independent glass transitions originating from the two components were  
173 clearly detected at almost the same positions as those observed for the unblended samples.  
174 This behaviour of double  $T_g$ s was also noted for  $CP_{2.81}/PVP$  and  $CP_{2.93}/PVP$  blends. Thus,  
175 the CPs of  $DS > 2.7$  are taken as immiscible with PVP.

## 176 <<Figure 2 (a) & (b)>>

177 Contrastively, the other six pairs of CP/PVP using propionyl DSs of  $<2.7$  imparted a  
178 miscible sign. Figure 2b exemplifies DSC thermograms of  $CP_{2.62}/PVP$  blends.  $T_g$  of  $CP_{2.62}$   
179 was determined to be  $138\text{ }^{\circ}\text{C}$ . The blends with PVP gave a single, composition-dependent  
180  $T_g$  that shifted to higher temperatures along with an increase in the PVP content; thus we can  
181 conclude that the CP forms a miscible monophase with PVP. This was also the case for the  
182 other CPs of  $DS = 1.71\text{--}2.54$ .

183 Figure 3 compiles FT-IR spectra obtained for blends of the miscible  $CP_{1.71}/PVP$  pair, on  
184 an enlarged scale for two regions of (a) O-H and (b) C=O stretching vibrations. As shown in  
185 Figure 3a, the unblended CP (top data) gave a band centering at  $3,482\text{ cm}^{-1}$ , which can be  
186 associated with a mixture of free hydroxyls and intramolecularly hydrogen-bonded OH  
187 groups. For the blends, it was observed that the band peak shifted to lower wavenumber  
188 positions with increasing PVP content, and, concomitantly, another absorption signal became

more discernible as a shoulder on the side of further lower wavenumbers, as marked by a white arrow at  $\sim 3,300\text{ cm}^{-1}$  in Figure 3a. This new band can be ascribed to the stretching of intermolecularly hydrogen-bonded OH groups (Marchessault and Liang 1960).

### <<Figure 3 (a) & (b)>>

Concerning the region of C=O stretching vibration (Fig. 3b), a  $1,744\text{ cm}^{-1}$  band involved in the propionyl side-group of the CP component was almost unchanged in the peak location by the blending with PVP. However, a carbonyl signal of PVP, observed at  $1,675\text{ cm}^{-1}$  for the homopolymer, became asymmetric progressively as the CP content increased in the binary mixture; consequently, the absorption band was dividable into two peaks, a larger one at  $\sim 1,680\text{ cm}^{-1}$  and a smaller one at  $\sim 1,660\text{ cm}^{-1}$  (see data for CP-rich compositions in Fig. 3b). These two split IR signals for the PVP component may be associated with the free carbonyl and hydrogen-bonded carbonyl groups, respectively (Masson and Manley 1991).

The above observations of the frequency shift and shape variation for the specific IR bands are evidently attributed to the hydrogen-bonding interaction between the residual hydroxyls of the CP component and the carbonyls of the PVP component. Conversely, this attractive interaction would contribute as a driving force to develop the good miscibility of the CP/PVP blends, as did in the CA/PVP (Miyashita et al. 2002; Ohno et al. 2005) and CB/PVP systems (Ohno and Nishio 2006). In a corroborating experiment, the immiscible CP/PVP blends using highly substituted CPs of DS = 2.81 and 2.93 exhibited no systematic variation of the corresponding bands in their FT-IR spectra.

By comprehensive comparison with the previous estimation shown in Figure 1, we notice that the upper limit in DS of CP miscible with PVP, which is  $\sim 2.7$ , is just intermediate between the corresponding ones, 2.8 and 2.5, for CA and CB, respectively. This is readily interpretable as due to the difference in effectiveness of the steric hindrance between the three ester side-groups which can inhibit the hydrogen-bonding interaction stated above, in consideration of the order of bulkiness for the acetyl, propionyl, and butyryl substituents.

215

## 216 *CP/PVAc blends*

217 As-cast films of CP/PVAc blends were mostly transparent to the naked eye (i.e., optically  
218 compatible) over the whole composition range. However, taking account of the refractive  
219 index 1.47–1.49 of CP, close to that of PVAc (1.4665 (Seferis 1999)), we should note that the  
220 transparency of these films is not directly linked to the blend miscibility.

221 Figure 4 collects  $T_g$  versus composition plots for eight series of CP/PVAc blends  
222 (propionyl DS = 1.90–2.93). As can be seen from the plots, the three blend series using CPs  
223 of DS = 1.90, 2.18, and 2.35 were completely immiscible, because two  $T_g$  signals appeared  
224 without any noticeable shift from their original locations for the two components. Regarding  
225 the other blend series using CPs of DS > 2.5, however, an appreciable extent of  $T_g$  shift was  
226 detected for both of the two components at compositions of CP/PVAc = 60/40–90/10,  
227 indicating that a certain amount of the CP constituent was dissolved into the PVAc phase, and  
228 *vice versa*. Therefore, we judge the CP(DS > 2.5)/PVAc pairs to be partially miscible.  
229 Such partial miscibility was never definable to the CA/PVAc and CB/PVAc systems  
230 irrespective of DS of the CA or CB component; any blend of both systems provided two  
231 invariable  $T_g$ s independent of the mixing composition.

232 <<Figure 4>>

233 The finding of the partial miscibility (or better compatibility) for the pairs of highly  
234 propionylated CP/PVAc is quite significant in the present study, as embodied below for CP  
235 blends with the copolymer P(VP-*co*-VAc). A structural affinity between the propionyl  
236 side-group (CH<sub>3</sub>-CH<sub>2</sub>-CO-O-C-) and the VAc unit (-(CH<sub>2</sub>-CH(-O-CO-CH<sub>3</sub>))-) might be  
237 responsible to the advent of the partial miscibility, as we have pointed out a similar effect in  
238 former studies on CE/poly( $\epsilon$ -caprolactone) blends (Nishio et al. 1997; Kusumi et al. 2008).  
239 The two structural unities containing a carbonyl moiety may be favorable for a relatively  
240 weak interaction of dipole-dipole antiparallel alignment. The presence of such a weak

interaction is also suggested in earlier papers dealing with a miscible system of PVAc with poly(methyl acrylate) ( $-(\text{CH}_2-\text{CH}(\text{COOCH}_3))_n-$ ) (Nandi et al. 1985; Takegoshi et al. 1993).

#### *CP/P(VP-co-VAc) blends*

In visual appearance, as-cast films of CP blends with VP-VAc copolymers were homogeneous and transparent, except for films of several polymer pairs composed of CP of DS > 2.7 and P(VP-co-VAc) having more than 70 mol% VP residues.

Figure 5 displays  $T_g$  variations with mixing composition for eight series of CP<sub>1.90</sub>/P(VP-co-VAc) blends, the VP fraction of the copolymer component ranging from 10 to 87 mol%. In the data plotting, when the VP fraction in P(VP-co-VAc) was  $\geq 23$  mol%, any blend series of CP<sub>1.90</sub>/P(VP-co-VAc) provided a smooth variation of a single  $T_g$  situated between the  $T_g$  values of the two unblended components. Thus, it turns out that CP<sub>1.90</sub> forms a miscible monophase with P(VP-co-VAc)s of VP > ~20 mol%. For selected blends of the miscible pairs, the presence of the hydrogen-bonding interaction between the CP-hydroxyl and VP-carbonyl groups was also ascertained by FT-IR measurements. With regard to a series of CP<sub>1.90</sub>/P(VP<sub>0.10</sub>-co-VAc<sub>0.90</sub>), a few samples of 60–80 wt% CP content gave two discrete  $T_g$ s, yet there occurred a noticeable extent of  $T_g$  shift over all the blend compositions, as can be seen from Figure 5. Therefore, exceptionally, this polymer pair is evaluated to be partially miscible.

#### <<Figure 5>>

In the same way, CP<sub>2.18</sub> showed a similar miscibility behaviour to that of CP<sub>1.90</sub>; viz., the CP was partially miscible with P(VP<sub>0.10</sub>-co-VAc<sub>0.90</sub>) and completely miscible with the other copolymers of VP:VAc = 23:77–87:13. Intriguingly, CP<sub>2.35</sub>, CP<sub>2.54</sub>, and CP<sub>2.62</sub> were all completely miscible even with P(VP<sub>0.10</sub>-co-VAc<sub>0.90</sub>) as well as with the others of VP > 20 mol%.

When the propionyl DS of the CP component reached 2.72 and more, the CPs were



miscible with P(VP-*co*-VAc)s of ca. 10–65 mol% VP residues, despite their imperfect miscibility with PVAc and PVP homopolymers. Figure 6 exemplifies the miscible evidence in DSC for CP<sub>2.89</sub>/P(VP<sub>0.52</sub>-*co*-VAc<sub>0.48</sub>) and CP<sub>2.89</sub>/P(VP<sub>0.10</sub>-*co*-VAc<sub>0.90</sub>) combinations. Accordingly, it follows that the CP/P(VP-*co*-VAc) system exhibited a definite miscibility window, as did the previous CB/P(VP-*co*-VAc) system (see Fig. 1b). As for the miscibility window of the latter system, it was reasonably concluded that a greater repulsion between the VP and VAc units in the random copolymer was mainly contributory to the miscibility attainment; this was rationalized by assessment of the Krigbaum-Wall interaction parameters (Ohno and Nishio 2007). The intramolecular copolymer effect may also be applicable to the present CP(DS > 2.7)/P(VP-*co*-VAc) blends.

#### <<Figure 6 (a) & (b)>>

It should be stressed here that the CPs of DS > 2.7 were miscible with P(VP<sub>0.10</sub>-*co*-VAc<sub>0.90</sub>) abundant in VAc residues, as was the case for the ones of DS = 2.35–2.62. At the comonomer ratio of VP:VAc = 10:90, the intramolecular repulsion effect would decline to a considerable extent; instead, however, the interaction coming from the structural affinity between the ester side-group of CP and the VAc unit of the copolymer would be more prevailing. It can therefore be assumed that, as a result of favorable balance of the two effects, the high-substituted CPs were miscible with P(VP<sub>0.10</sub>-*co*-VAc<sub>0.90</sub>). To find a spectroscopic evidence of the latter interaction in which both CP- and VAc-carbonyls should be involved, we carried out FT-IR and solid-state <sup>13</sup>C CP/MAS NMR measurements for CP<sub>2.89</sub>/P(VP<sub>0.10</sub>-*co*-VAc<sub>0.90</sub>) and CP<sub>2.89</sub>/P(VP<sub>0.52</sub>-*co*-VAc<sub>0.48</sub>) blends. In the IR examination, however, the C=O stretching band of CP overlapped completely with the one of VAc. In the CP-MAS spectra measurements, the unblended copolymers gave a carbonyl resonance signal composed of two splitting peaks with their maximum at 171 ppm (for VAc unit) and 175 ppm (for VP unit); the splitting was relatively clear in a data for P(VP<sub>0.52</sub>-*co*-VAc<sub>0.48</sub>) (see Fig. 9). Nevertheless, when the carbonyl carbon resonance of CP (173.5 ppm) merged with the split

signal of the VP/VAc units, it was difficult to precisely estimate the respective three chemical shifts. Thus the structural affinity effect was undetectable for any of the blends; it appears to be substantially feeble, however.

On the basis of the thermal analysis data, we successfully constructed a miscibility map for the CP/P(VP-*co*-VAc) system, as shown in Figure 7. The diagram indicates that CPs of  $DS < 2.7$ , having a relatively higher amount of residual OH groups, are mostly miscible with the vinyl polymers of VP  $> \sim 20$  mol%, primarily due to predominance of the hydrogen-bonding interaction. A miscibility window emerges in the region satisfying propionyl  $DS > 2.7$  and VP fraction = 10–65 mol%, as a result of indirect intercomponent attraction due to the stronger repulsion effect in the P(VP-*co*-VAc) component itself. To make a comparison of the map with the previous ones (Fig. 1) for the corresponding blend systems of CA and CB, we find that the CP system produced the largest miscible region. As compared with the map for the CB blends, the DS boundary partitioning the miscibility states of CP/P(VP-*co*-VAc) (VP  $\geq 65$  mol%) pairs is driven up to  $\sim 2.7$  from the value  $\sim 2.5$  for the CB system. This elevation in DS may be ascribed to the modest effectiveness in steric hindrance of the propionyl group of medium size, relative to that of the more bulky butyryl group. Another factor expanding the miscible region in the map for the CP system is an intermolecular accessibility derived from the structural affinity of the propionyl side-group with the VAc unit; this effect may be applicable to the blending pairs of CPs of  $DS > \sim 2.3$  and P(VP-*co*-VAc)s of VP = ca. 10–20 mol%.

<<Figure 7>>

Insight into the scale of homogeneous mixing

In the previous study on the CB/P(VP-*co*-VAc) system (Ohno and Nishio 2006), it was suggested that the degree of homogeneity for the combinations situated in the miscibility

window was somewhat lower than that for the hydrogen-bonding type of miscible blends, reflecting a difference in absolute strength between the driving forces for the respective miscibility attainments. In this connection, we should remark that DSC thermograms for the 60/40–90/10 compositions of CP<sub>2.89</sub>/P(VP<sub>0.10</sub>-co-VAc<sub>0.90</sub>) blends exhibited a single, but relatively broader glass transition (see Fig. 6b). A similar phenomenon was noted for other CP/P(VP-co-VAc) blends satisfying  $DS > 2.7$  for the CP and VP = 10–33 mol% for the copolymer. Such a broadening in temperature range of the glass transition may be interpretable as due to mixing of plural microdomains with subtly different fluctuations in polymer composition (Lodge and McLeish 2000). In relation to this assumption, further investigations were made into the homogeneity of mixing for the CP/P(VP-co-VAc) system by means of DMA and nuclear magnetic relaxation measurements.

#### *Thermal transition behavior evaluated by DMA*

As far as detection of  $T_g$  is concerned, DMA is more sensitive than calorimetric measurements in many cases of studies on multicomponent polymeric materials (Kusumi et al. 2011; MacKnight et al. 1978; Ultracki 1990). As a conventional matter, if DSC analysis is sensitive to heterogeneities with sizes of ca. 20–30 nm as an upper limit, DMA can detect a somewhat finer scale of heterogeneity, e.g., a domain size smaller than ~15 nm (Kaplan 1976; Masson and Manley 1991; Nishio 1994).

Figure 8a shows the temperature dependence of the dynamic storage modulus  $E'$  and loss modulus  $E''$  for CP<sub>2.89</sub>/P(VP<sub>0.10</sub>-co-VAc<sub>0.90</sub>) blends of 25/75, 50/50, and 75/25 compositions, together with the corresponding data for plain CP<sub>2.89</sub>. As for the copolymer *per se*, the data was not obtained because of a brittle nature of the film. As demonstrated clearly in the figure, the unblended CP<sub>2.89</sub> sample showed a very sharp transition with an  $E''$  peak maximum at 137 °C; this temperature is somewhat higher than  $T_g$  (127 °C) determined by DSC, however. In contrast, the blend samples gave a much broader  $E''$  peak with a low

onset point (~50 °C in common), and a more gradual  $E'$  falling as well, in the glass transition temperature region, this trend being particularly prominent in the data for the 75/25 and 50/50 compositions.

### <<Figure 8 (a) & (b)>>

Figure 8b compiles DMA data of CP<sub>2.18</sub>/PVP and CP<sub>2.89</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) blends, the illustration for each series being restricted to a few samples rich in CP content, there. Any of the two blend series provided a single and sharp transition signal, both in the  $E''$  peak and in the  $E'$  drop, which shifted systematically with polymer composition. From comparison with these data, the pair of CP<sub>2.89</sub>/P(VP<sub>0.10-co</sub>-VAc<sub>0.90</sub>) described on ahead is obviously inferior in the degree of miscibility to the other two, within a scale (~15 nm) detectable by DMA. Incidentally, it is interesting to find a homogeneity on the DMA scale for the CP<sub>2.89</sub> blends with P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) having equimolar amounts of VP and VAc units, although the blending pair is not of the hydrogen-bonding type such as the CP<sub>2.18</sub>/PVP one but situated in the miscibility window of that map (Fig. 7).

### *Homogeneity estimated by solid-state <sup>13</sup>C NMR relaxation*

As a useful technique in solid-state <sup>13</sup>C NMR,  $T_{1\rho}^H$  measurements for specific carbons in a multicomponent polymer system make it possible to estimate the mixing homogeneity in a scale of <sup>1</sup>H spin-diffusion length that is usually within several nanometers (Masson and Manley 1991; Ohno et al. 2005; Zhang et al. 1992).  $T_{1\rho}^H$  values can be obtained by fitting the decaying carbon resonance intensity to the following exponential equation:

$$M(\tau) = M(0) \exp(-\tau/T_{1\rho}^H) \quad (1)$$

where  $M(\tau)$  is the magnetization intensity observed as a function of the spin-locking time  $\tau$ . In a general rule, if two constituent polymers are in a homogeneously mixed state on the scale over which <sup>1</sup>H spin-diffusion can take place in a time  $T_{1\rho}^H$ , the  $T_{1\rho}^H$  values for different protons belonging to the respective components may be equalized to each other by the spin

diffusion.

In terms of the NMR technique, a comparative assessment of the polymer-polymer mixing scale was made for three selected series of blends, CP<sub>2.89</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), CP<sub>1.71</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), and CP<sub>1.71</sub>/PVP. Figure 9 exemplifies <sup>13</sup>C CP/MAS spectra obtained for CP<sub>2.89</sub>, P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), and their 50/50 blend. The peak assignments of the spectra are based on literature data for CP (Tezuka and Tsuchiya 1995), PVP (Zhang et al. 1992), and PVAc (Cheung et al. 2000). The experiment of  $T_{1\rho}^H$  quantifications was done through monitoring the following <sup>13</sup>C resonance signals with better resolution: C2/C3/C5 pyranose carbons (74 ppm) and propionyl carbons C8 (28 ppm) and C9 (9.3 ppm) for the CP component, and C<sub>b</sub>/C<sub>c</sub> (42 ppm) and C<sub>δ</sub>/C<sub>d</sub> carbons (~20 ppm) for the P(VP-co-VAc) component. Figure 10a illustrates the decay behaviour in intensity of the C2/C3/C5 and C<sub>b</sub>/C<sub>c</sub> peaks for unblended CP<sub>2.89</sub> and P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), respectively, and for their 50/50 blend imparting both resonance signals as well. The slope of each semi-logarithmic plot corresponds to an inverse of  $T_{1\rho}^H$  as the time constant of the relaxation process. We found from these plots that  $T_{1\rho}^H$  of CP<sub>2.89</sub> (18.5 ms) increased to 22.7 ms and that of P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) (35.4 ms) decreased to 33.1 ms by the 50/50 blending, but they never became so close to each other. Regarding the CP<sub>1.71</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) pair (Fig. 10b), on the contrary,  $T_{1\rho}^H$  values of the two components for the 50/50 blend coincided with each other just at the midpoint (27.9 ms) between the respective original values, 20.5 ms for CP<sub>1.71</sub> and 35.4 ms for the copolymer. A similar tendency of  $T_{1\rho}^H$  variations was observed in tracing of another set of the C8 or C9 signal of CP and the C<sub>δ</sub>/C<sub>d</sub> signal of P(VP-co-VAc).

<<Figure 9>>

<<Figure 10 (a) & (b)>>

Table 2 lists all the  $T_{1\rho}^H$  data obtained for CP<sub>1.71</sub>, CP<sub>2.89</sub>, PVP, P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), and their miscible blends of CP/vinyl polymer = 75/25–25/75. In the CP<sub>1.71</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) series,  $T_{1\rho}^H$  of the CP<sub>1.71</sub> component, originally 20.0 ms as an average, rises systematically

with an increase in the copolymer content, while that of the P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) component, originally 35.0 ms as an average, diminishes correspondingly with increasing CP<sub>1.71</sub> content. In consequence, the two  $T_{1\rho}^H$ s at every blend composition are surely in good agreement with each other. Such a composition-dependent shift of the almost equalized  $T_{1\rho}^H$ s of two components is also observed for the CP<sub>1.71</sub>/PVP blend series. Thus, it is reasonably deduced that the two constituent polymers in the two series of blends are intimately mixed within a range where the mutual <sup>1</sup>H-spin diffusion is permitted over a period of the respective homogenized  $T_{1\rho}^H$ , e.g., ~30.5 ms for the 50/50 composition of CP<sub>1.71</sub>/PVP.

## <<Table 2>>

An effective path length  $L$  of the spin diffusion in a time  $T_{1\rho}^H$  is given by the following equation (McBrierty and Douglass 1981):

$$L \cong (6DT_{1\rho}^H)^{1/2} \quad (2)$$

where  $D$  is the spin-diffusion coefficient, usually taken to be  $\sim 1.0 \times 10^{-12}$  cm<sup>2</sup>/s in organic polymer materials. By adopting  $T_{1\rho}^H$  data of 23–31 ms approximated for the CP<sub>1.71</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) blends of 75/25–25/75 compositions, the diffusion path length is calculated as  $L = 3.7$ – $4.3$  nm. In a similar manner,  $L$  is determined to be  $3.9$ – $4.4$  nm with  $T_{1\rho}^H = 25$ – $33$  ms for the corresponding CP<sub>1.71</sub>/PVP compositions. Accordingly, it is confirmed that any of these miscible blends using a low-substituted CP is virtually homogeneous in a scale of ca. 4 nm.

With regard to the CP<sub>2.89</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) series, as evidenced in Table 2, there arises a serious disagreement between  $T_{1\rho}^H$ s of the two polymer components at every blend composition, although the mutual approach to a small extent is admitted. This larger temporal disagreement implies that the relaxation processes of the two polymers in the blends proceeded rather independently without their cooperative spin diffusion. By the combined use of this result and the previous DMA one, it can be concluded that the scale of homogeneity in the CP<sub>2.89</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) blends lies between approximately 5 and 15

423 nm.

424

## 425 Conclusions

426

427 Miscibility characterization was performed on blends of cellulose propionate (CP) with  
428 synthetic vinyl polymers containing *N*-vinyl pyrrolidone (VP) and/or vinyl acetate (VAc) units,  
429 i.e., PVP, PVAc, and P(VP-*co*-VAc) random copolymers. On the basis of  $T_g$  analysis by  
430 DSC, a miscibility map (Fig. 7) was successfully constructed as a function of both the  
431 propionyl DS of CP and the VP:VAc composition of P(VP-*co*-VAc). FT-IR spectroscopy  
432 was also utilized to detect a hydrogen-bonding type of intermolecular interaction contributory  
433 to the miscibility attainment. As denoted in that map, polymer pairs of CP/P(VP-*co*-VAc)  
434 satisfying  $DS < 2.7$  for the CP component and  $VP > 20$  mol% for the vinyl polymer  
435 component were miscible. This miscibility is given rise to, more or less, by virtue of the  
436 hydrogen bonding between CP-hydroxyls and VP-carbonyls, and hence the effectiveness  
437 should be greater when the propionyl DS is lower and the VP fraction is higher. The upper  
438 limit of  $DS = 2.7$  required for the miscibility was intermediate between the corresponding  
439 ones, acetyl  $DS = 2.8$  and butyryl  $DS = 2.5$ , for the comparable systems employing cellulose  
440 acetate (CA) and butyrate (CB). This observation can be explained as being due to the  
441 difference in bulkiness between the three sorts of acyl substituents, each exerting an effect of  
442 steric hindrance to decline the hydrogen-bonding interaction.

443 CPs of  $DS > 2.7$  exhibited miscibility with P(VP-*co*-VAc)s of  $VP = 10$ – $65$  mol%,  
444 despite their imperfect miscibility with both PVP and PVAc homopolymers; this resulted in  
445 advent of a definite 'miscibility window' in the map, as has been experienced formerly in our  
446 study of CB/P(VP-*co*-VAc) blends. The behaviour may be interpreted to be occasioned  
447 principally by intramolecular repulsion between the comonomer units in P(VP-*co*-VAc). In  
448 addition, another effect, the structural affinity between the propionyl side-group and the VAc



unit, contributes to the miscibility realized for blends of highly propionylated CPs ( $DS > 2.3$ ) with VAc-rich copolymers (VP:VAc = 10:90–33:67). Eventually, the miscible pairing region extended more widely in the map for the CP/P(VP-*co*-VAc) system, compared with the situations in the corresponding blend systems using CA and CB. It is astonishing afresh to find that only one difference in carbon number of the acyl substitution drastically changed the miscibility behaviour of cellulose esters (CEs) with a given synthetic copolymer.

However, caution should be exercised to the scale of homogeneity in the CP/P(VP-*co*-VAc) blends being estimated to form a miscible monophasic. As a result of further investigation by DMA and  $T_{1\rho}^H$  quantifications in solid-state  $^{13}\text{C}$  NMR, we had awareness of the following respects: The miscible blends of hydrogen-bonding type are completely homogeneous on a scale of a few nanometers ( $\leq 4$  nm), whereas the blend series situated in the miscibility window are homogeneous with a possible microdomain size between ca. 5 and 15 nm. In the latter assessment, CP blends with P(VP-*co*-VAc)s extremely rich in VAc (e.g., VP:VAc = 10:90) are excepted from the relevant group, their miscibility being invited by a rather weak interaction due to the structural affinity effect. The blends of exception can contain heterogeneous domains of ca. 15–20 nm sizes as a tentative estimate.

From a practical standpoint, these results will contribute toward expanding the opportunities of material design based on the CE family. Further studies along this line of fundamental characterization are now in progress for CE blends with other vinyl copolymers, in parallel with inquiries into their functionalities as optical and/or membrane materials. In the not too distant future, our effort will also be made to investigate an effect of regioselectivity of the acyl substituent in CEs on their miscibility with synthetic polymers, beyond the discussion in terms of the average DS parameter.

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## Figure Captions

**Fig. 1** Miscibility maps for two blend systems (a) CA/P(VP-*co*-VAc) and (b) CB/P(VP-*co*-VAc), quoted from previous papers (Miyashita et al. 2002; Ohno and Nishio 2006) in a rearranged style retaining the essence.

**Fig. 2** DSC thermograms obtained for (a) CP<sub>2.72</sub>/PVP and (b) CP<sub>2.62</sub>/PVP blends. Arrows indicate a  $T_g$  position taken as the midpoint of a baseline shift in heat flow.

**Fig. 3** FT-IR spectra of CP<sub>1.71</sub>, PVP, and their blends in the frequency regions of (a) O-H and (b) C=O stretching vibrations. Solid arrows indicate a peak-top position in the respective specific absorption bands, and white arrows indicate a shoulder band associated with hydrogen bonding (see text for discussion).

**Fig. 4**  $T_g$  versus composition plots for eight series of CP/PVAc blends. DS of CP: ○, 1.90; ●, 2.18; □, 2.35; ■, 2.54; △, 2.62; ▲, 2.72; ◇, 2.81; ◆, 2.93.

**Fig. 5** Composition dependence of  $T_g$  for eight series of CP<sub>1.90</sub>/P(VP-*co*-VAc) blends. VP:VAc in P(VP-*co*-VAc): ●, 10:90; ■, 23:77; ▽, 33:67; ◇, 40:60; +, 52:48; □, 62:38; △, 73:27; ○, 87:13.

**Fig. 6** DSC thermograms obtained for (a) CP<sub>2.89</sub>/P(VP<sub>0.52</sub>-*co*-VAc<sub>0.48</sub>) and (b) CP<sub>2.89</sub>/P(VP<sub>0.10</sub>-*co*-VAc<sub>0.90</sub>) blends. Arrows indicate a  $T_g$  position.

**Fig. 7** Miscibility map for CP/P(VP-*co*-VAc) blends, as a function of DS of CP and VP fraction in P(VP-*co*-VAc). Symbols indicate that a given pair of CP/P(VP-*co*-VAc) is

563 miscible (○), immiscible (×), or partially miscible (△).

564

565 **Fig. 8** Temperature dependence of the dynamic storage modulus  $E'$  and loss modulus  $E''$  for

566 (a) CP<sub>2.89</sub>/P(VP<sub>0.10-co</sub>-VAc<sub>0.90</sub>) and (b) CP<sub>2.18</sub>/PVP and CP<sub>2.89</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>) blends.

567

568 **Fig. 9** Solid-state  $^{13}\text{C}$  CP/MAS NMR spectra for CP<sub>2.89</sub>, P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), and their

569 50/50 blend.

570

571 **Fig. 10** Semilogarithmic plots of the decay of  $^{13}\text{C}$  resonance intensities as a function of

572 spin-locking time  $\tau$ , for solid films of (a) CP<sub>2.89</sub>, P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), and their 50/50 blend,

573 and (b) CP<sub>1.71</sub>, P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), and their 50/50 blend. The monitoring was conducted

574 for the peak intensity of C2/C3/C5 pyranose carbons of CP and that of C<sub>b</sub>/C<sub>c</sub> carbons of the

575 copolymer (see Fig. 9).

576

577 -----

578 In addition to the ten figures, there are two tables. See annexed sheets.

579 **Table 1** Characterization of CP and synthetic vinyl polymers used in the present study

Sample	DS <sup>a</sup>	$M_w$ <sup>b</sup>	$M_n$ <sup>b</sup>	$M_w/M_n$ <sup>b</sup>	$T_g / ^\circ\text{C}$	Source
CP	1.71	2,010,000	850,000	2.36	162	Synthesized
	1.90	1,860,000	824,000	2.26	161	Synthesized
	2.18	1,300,000	577,000	2.25	157	Synthesized
	2.35	2,210,000	925,000	2.39	153	Synthesized
	2.54	1,180,000	509,000	2.32	140	Synthesized
	2.62	979,000	359,000	2.73	138	Synthesized
	2.72	2,390,000	968,000	2.47	134	Synthesized
	2.81	1,990,000	837,000	2.38	128	Synthesized
	2.89	2,000,000	692,000	2.89	127	Synthesized
	2.93	1,250,000	525,000	2.38	124	Synthesized
Sample	VP content / mol% <sup>a</sup>	$M_w$ <sup>c</sup>	$M_n$ <sup>c</sup>	$M_w/M_n$ <sup>c</sup>	$T_g / ^\circ\text{C}$	Source
PVP	100	360,000 <sup>d</sup>	—	—	177	Nacalai Tesque, Inc.
P(VP-co-VAc)	87	56,500	28,000	2.02	124	Synthesized <sup>e</sup>
	73	52,100	25,000	2.08	111	Synthesized <sup>e</sup>
	62	51,600	24,400	2.11	101	Synthesized <sup>e</sup>
	52	28,000	5,120	5.47	89	Polyscience, Inc. <sup>f</sup>
	40	51,100	20,700	2.47	76	Synthesized <sup>e</sup>
	33	23,300	3,800	6.12	72	Polyscience, Inc. <sup>f</sup>
	23	59,400	26,100	2.27	55	Synthesized <sup>e</sup>
	10	46,500	27,700	1.68	45	Synthesized <sup>e</sup>
PVAc	0	90,000 <sup>d</sup>	—	—	41	Polyscience, Inc.

<sup>a</sup> Determined by <sup>1</sup>H NMR.

<sup>b</sup> Determined by gel permeation chromatography (mobile phase, tetrahydrofuran at 40 °C) with polystyrene standards.

<sup>c</sup> Determined by gel permeation chromatography (mobile phase, 10 mM/L lithium bromide/*N,N*-dimethylformamide at 40 °C) with polystyrene standards.

<sup>d</sup> Nominal value.

<sup>e</sup> Synthesized in our laboratory by radical polymerization of two distilled monomers, VP (Aldrich Chemical Co.) and VAc (Nacalai Tesque, Inc.), in the same way as that described in a previous paper (Miyashita et al. 2002).

<sup>f</sup> Used after purification by dissolution in dichloromethane and reprecipitation into petroleum ether.

581 **Table 2**  $T_{1\rho}^H$  values obtained for three series of blends, CP<sub>2.89</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>),  
582 CP<sub>1.71</sub>/P(VP<sub>0.52-co</sub>-VAc<sub>0.48</sub>), and CP<sub>1.71</sub>/PVP

CP <sub>2.89</sub> /P(VP <sub>0.52-co</sub> -VAc <sub>0.48</sub> ) (wt/wt)	$T_{1\rho}^H$ / ms						
	CP <sub>2.89</sub>				P(VP <sub>0.52-co</sub> -VAc <sub>0.48</sub> )		
	C2/3/5	C8	C9	Ave.	b/c	δ/d	Ave.
100/0	18.5	17.6	17.7	17.9	—	—	—
75/25	20.3	18.8	18.0	19.0	32.9	29.7	31.3
50/50	22.7	20.5	19.2	20.8	33.1	30.9	32.0
25/75	25.3	22.6	21.2	23.0	34.7	33.8	34.3
0/100	—	—	—	—	35.4	34.6	35.0

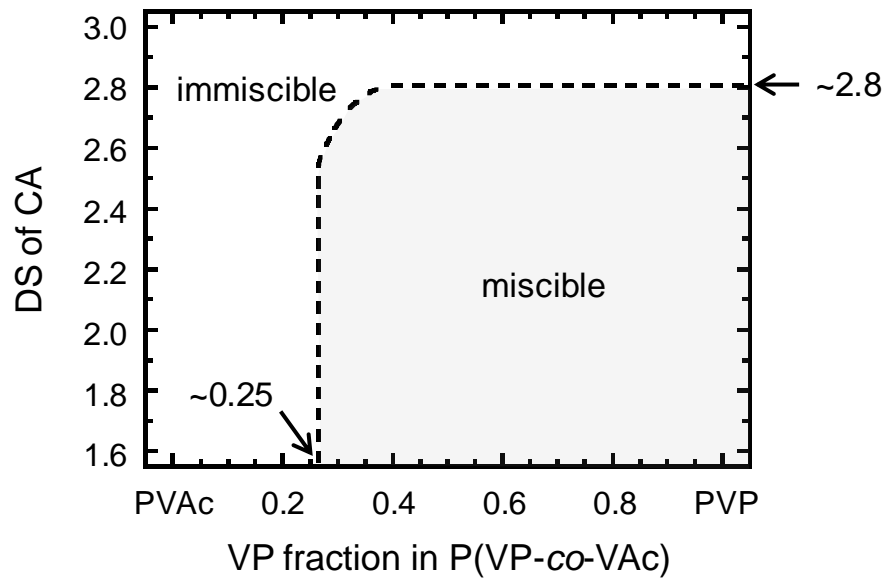
CP <sub>1.71</sub> /P(VP <sub>0.52-co</sub> -VAc <sub>0.48</sub> ) (wt/wt)	$T_{1\rho}^H$ / ms						
	CP <sub>1.71</sub>				P(VP <sub>0.52-co</sub> -VAc <sub>0.48</sub> )		
	C2/3/5	C8	C9	Ave.	b/c	δ/d	Ave.
100/0	20.5	20.3	19.2	20.0	—	—	—
75/25	24.3	23.0	21.6	23.0	24.5	24.1	24.3
50/50	27.8	26.8	24.5	26.4	27.9	26.9	27.4
25/75	33.0	29.7	27.8	30.2	33.7	30.9	32.3
0/100	—	—	—	—	35.4	34.6	35.0

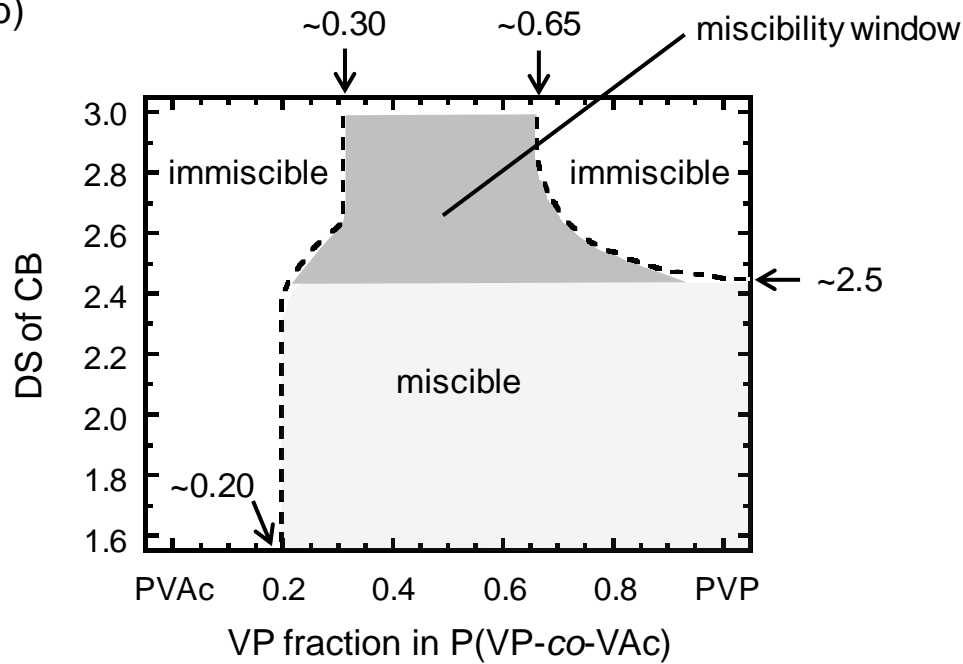
CP <sub>1.71</sub> /PVP (wt/wt)	$T_{1\rho}^H$ / ms						
	CP <sub>1.71</sub>				PVP		
	C2/3/5	C8	C9	Ave.	b/c	d	Ave.
100/0	20.5	20.3	19.2	20.0	—	—	—
75/25	25.1	24.1	23.0	24.1	26.0	26.0	26.0
50/50	30.9	30.1	30.8	30.6	30.6	29.5	30.1
25/75	33.4	33.7	33.1	33.4	33.7	33.0	33.4
0/100	—	—	—	—	32.6	31.3	32.0

583

(a)

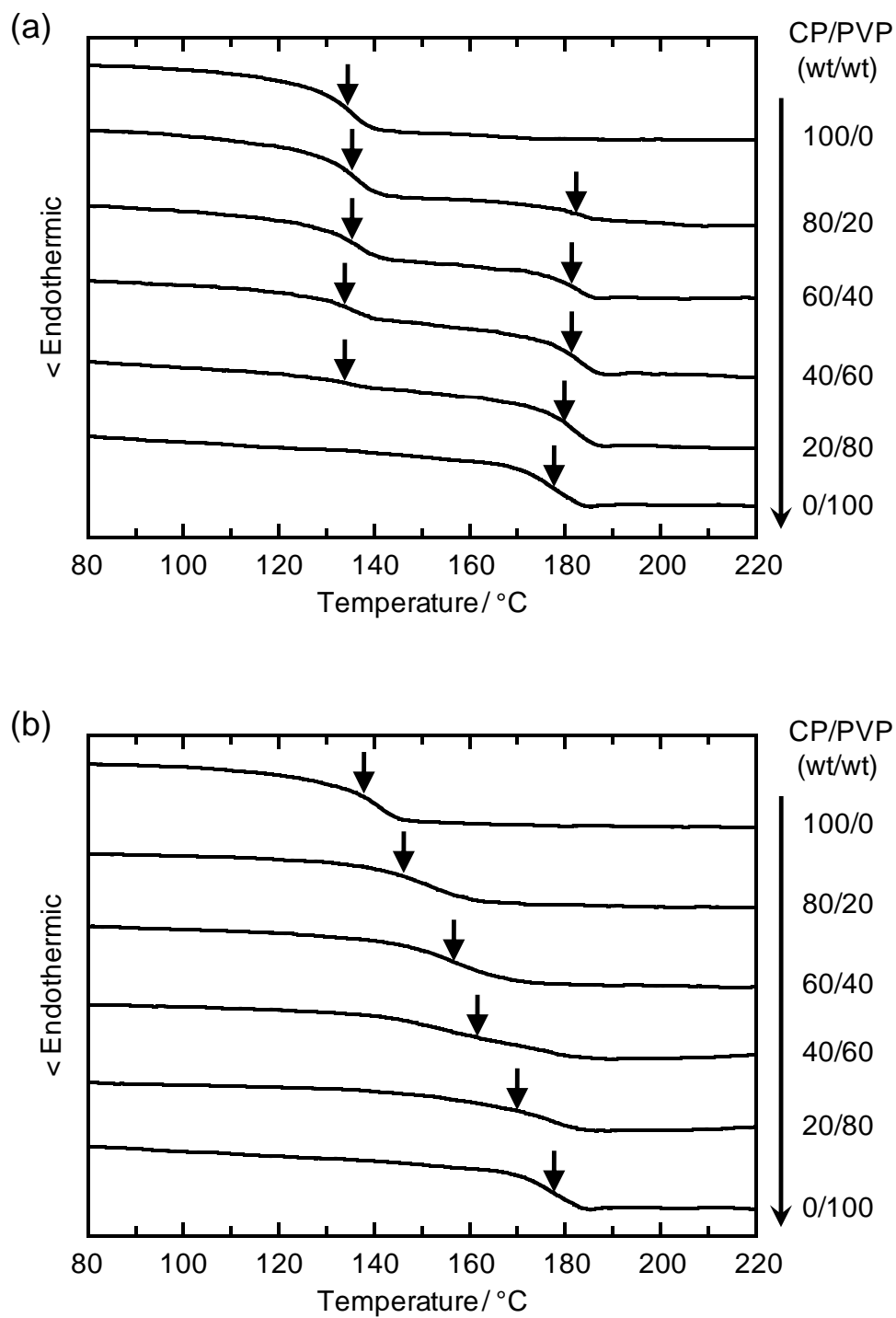


(b)

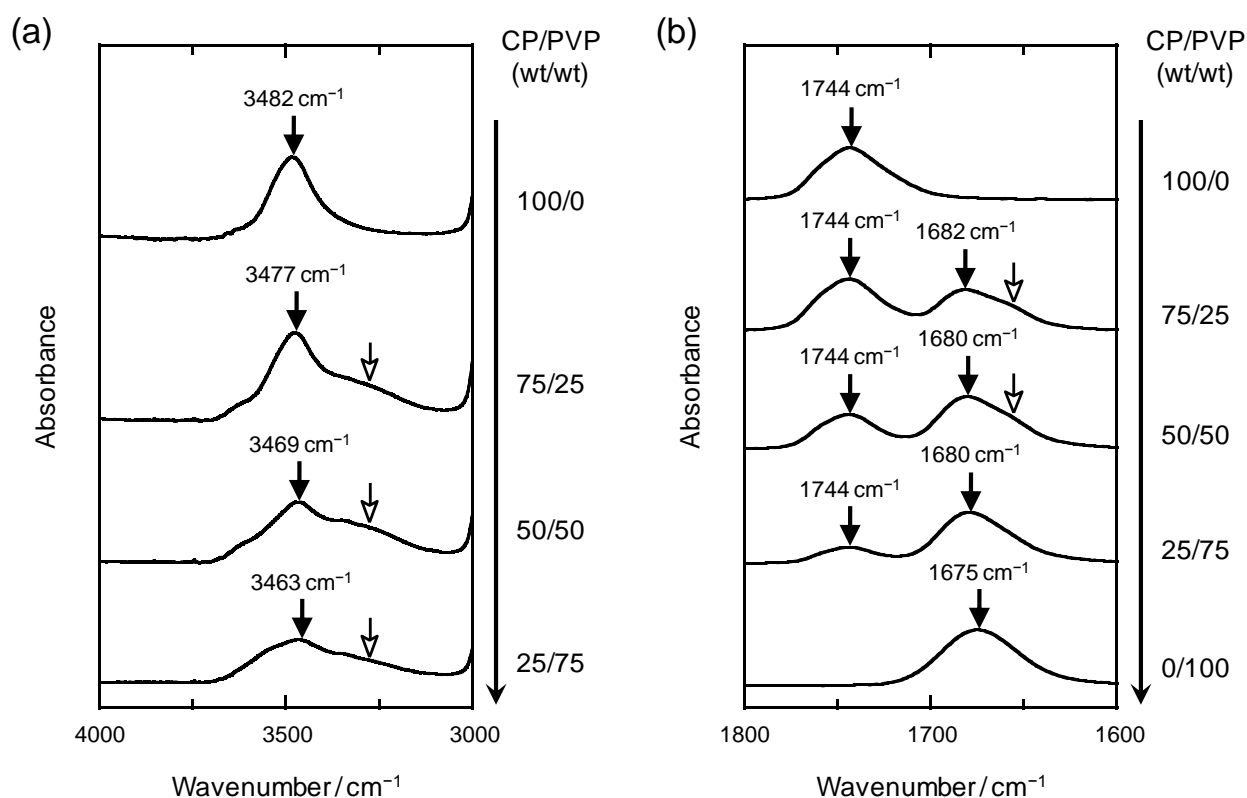


**Fig. 1** Miscibility maps for two blend systems (a) CA/P(VP-co-VAc) and (b) CB/P(VP-co-VAc), quoted from previous papers (Miyashita et al. 2002; Ohno and Nishio 2006) in a rearranged style retaining the essence.

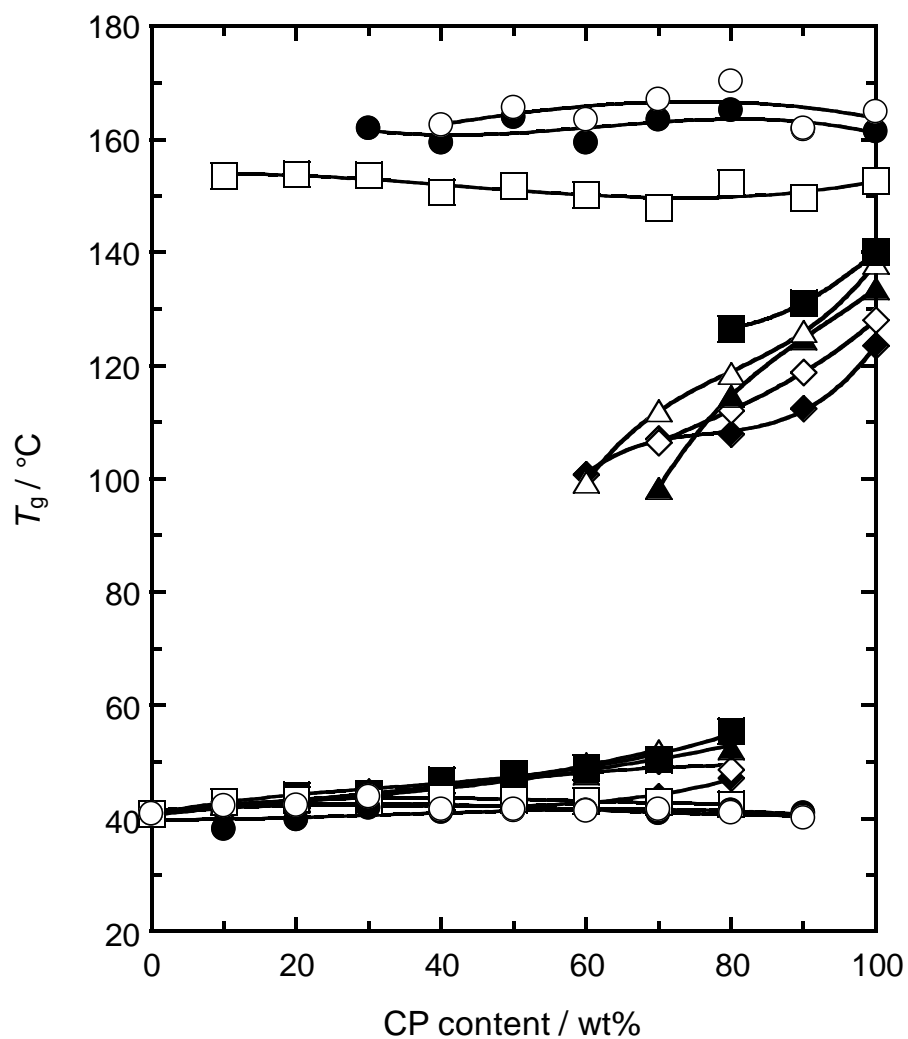




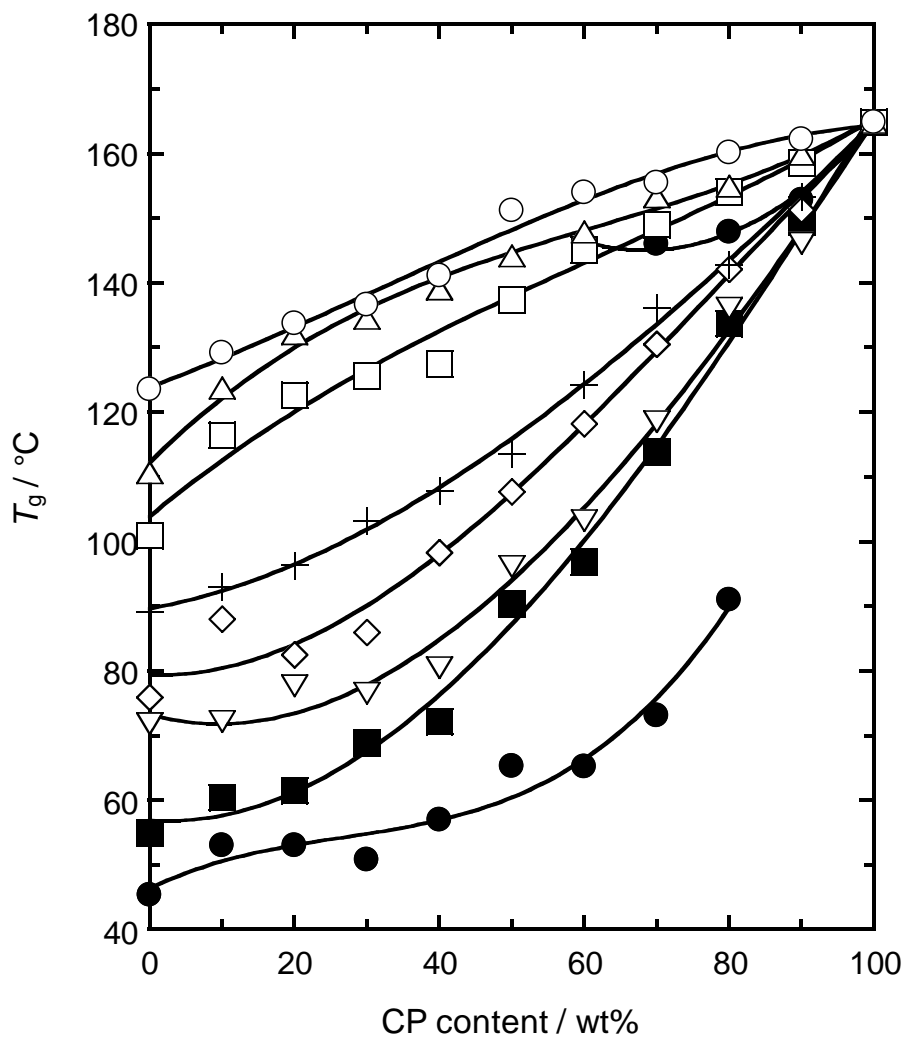
**Fig. 2** DSC thermograms obtained for (a) CP<sub>2.72</sub>/PVP and (b) CP<sub>2.62</sub>/PVP blends. Arrows indicate a  $T_g$  position taken as the midpoint of a baseline shift in heat flow.



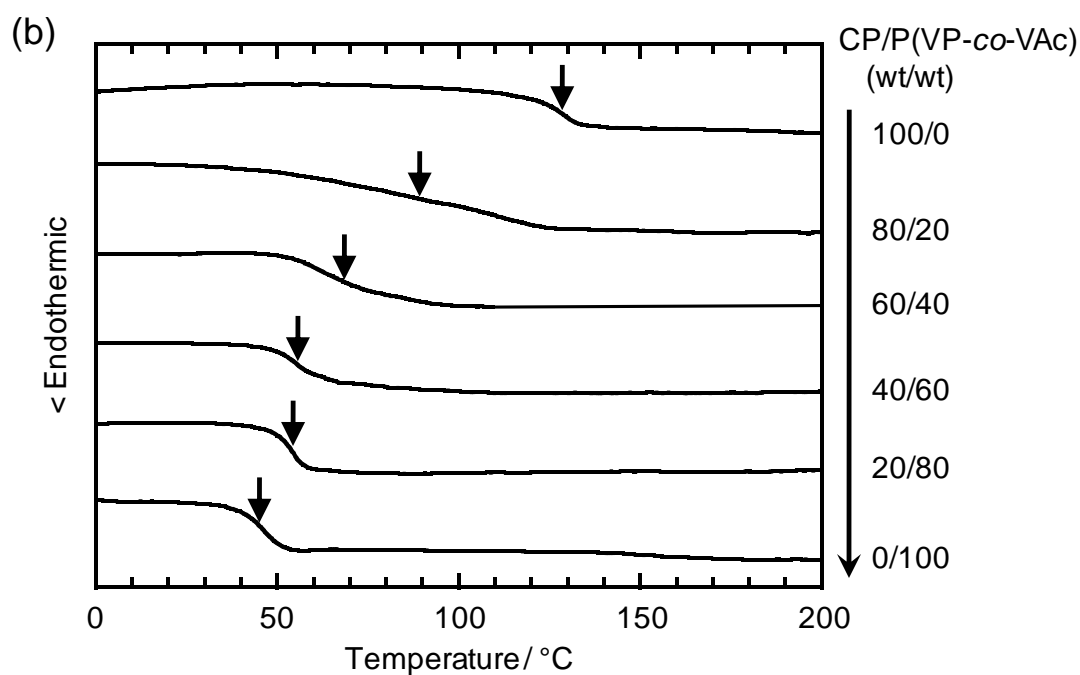
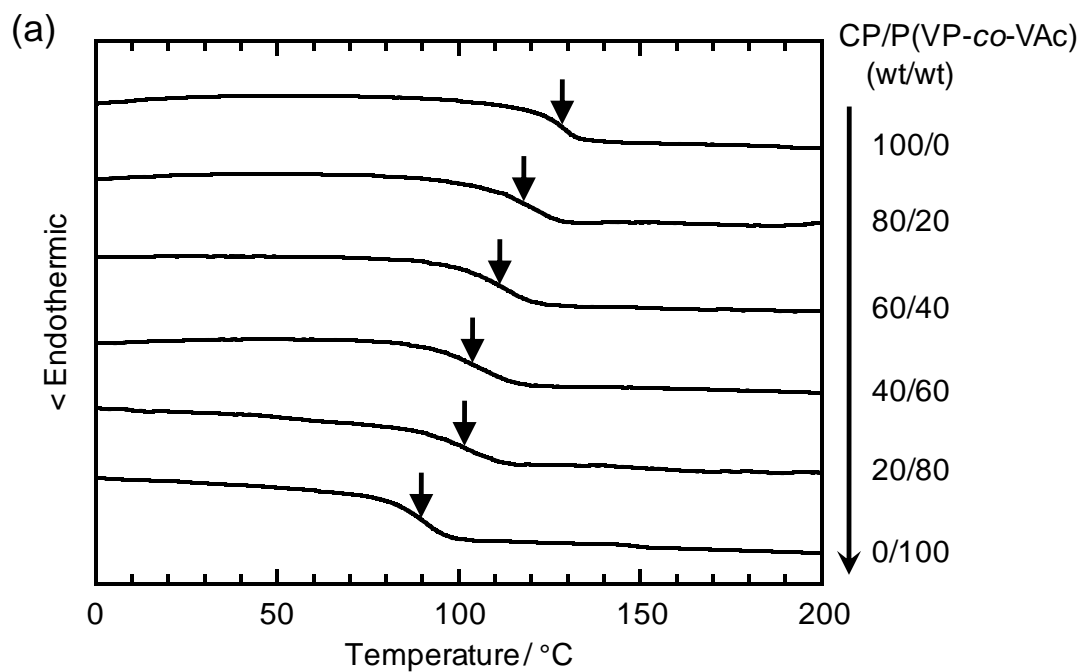
**Fig. 3** FT-IR spectra of CP<sub>1.71</sub>, PVP, and their blends in the frequency regions of (a) O-H and (b) C=O stretching vibrations. Solid arrows indicate a peak-top position in the respective specific absorption bands, and white arrows indicate a shoulder band associated with hydrogen bonding (see text for discussion).



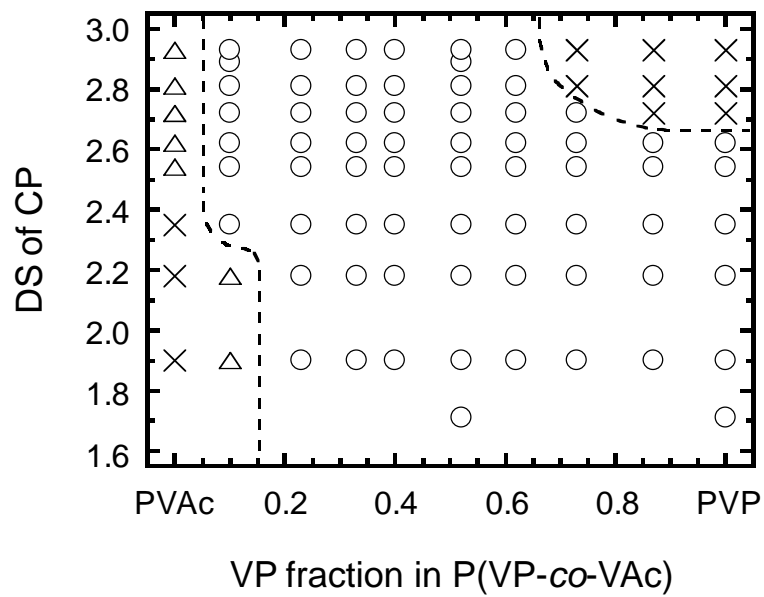
**Fig. 4**  $T_g$  versus composition plots for eight series of CP/PVAc blends. DS of CP: ○, 1.90; ●, 2.18; □, 2.35; ■, 2.54; △, 2.62; ▲, 2.72; ◇, 2.81; ◆, 2.93.



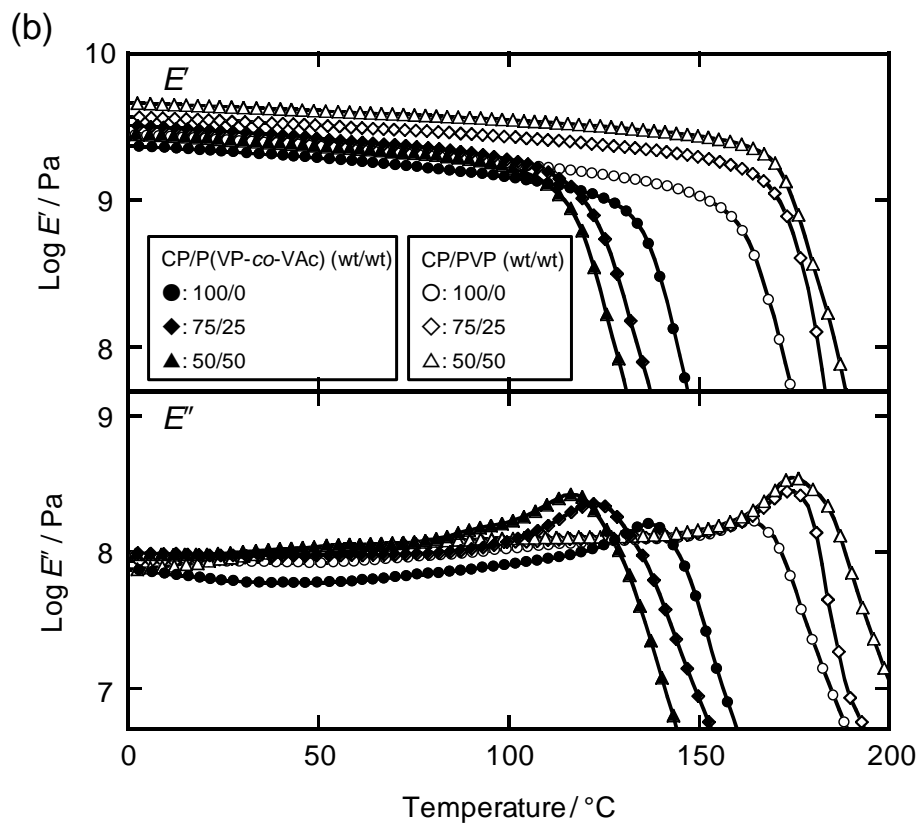
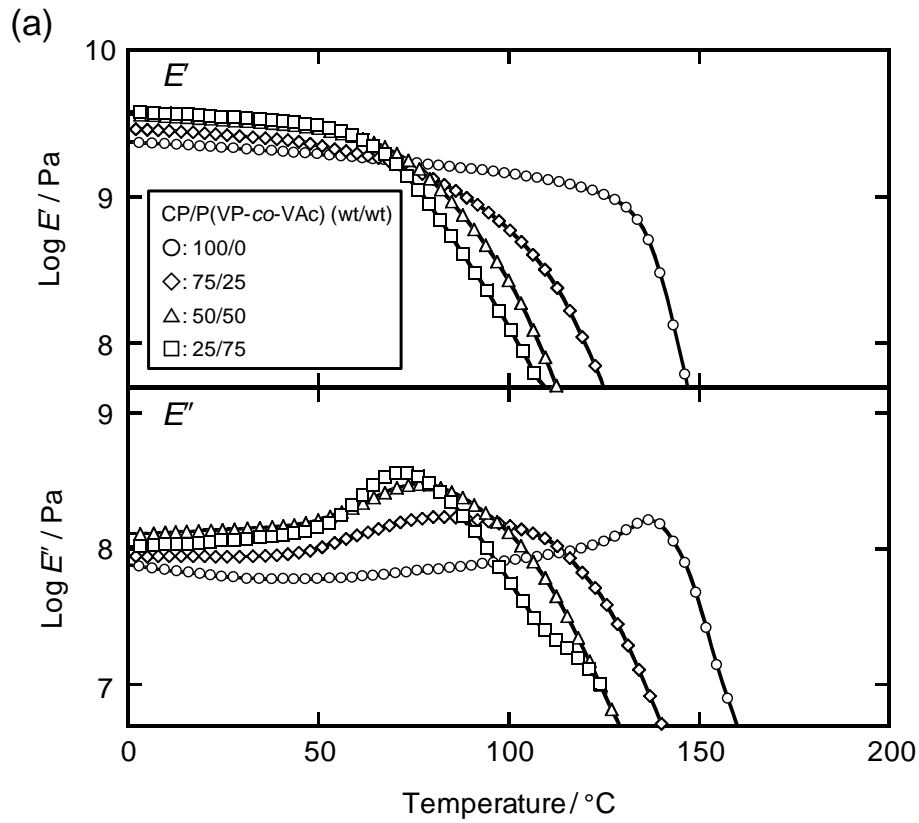
**Fig. 5** Composition dependence of  $T_g$  for eight series of  $CP_{1.90}/P(VP-co-VAc)$  blends.  
VP:VAc in  $P(VP-co-VAc)$ : ●, 10:90; ■, 23:77; ▽, 33:67; ◇, 40:60; +, 52:48; □, 62:38; △, 73:27; ○, 87:13.



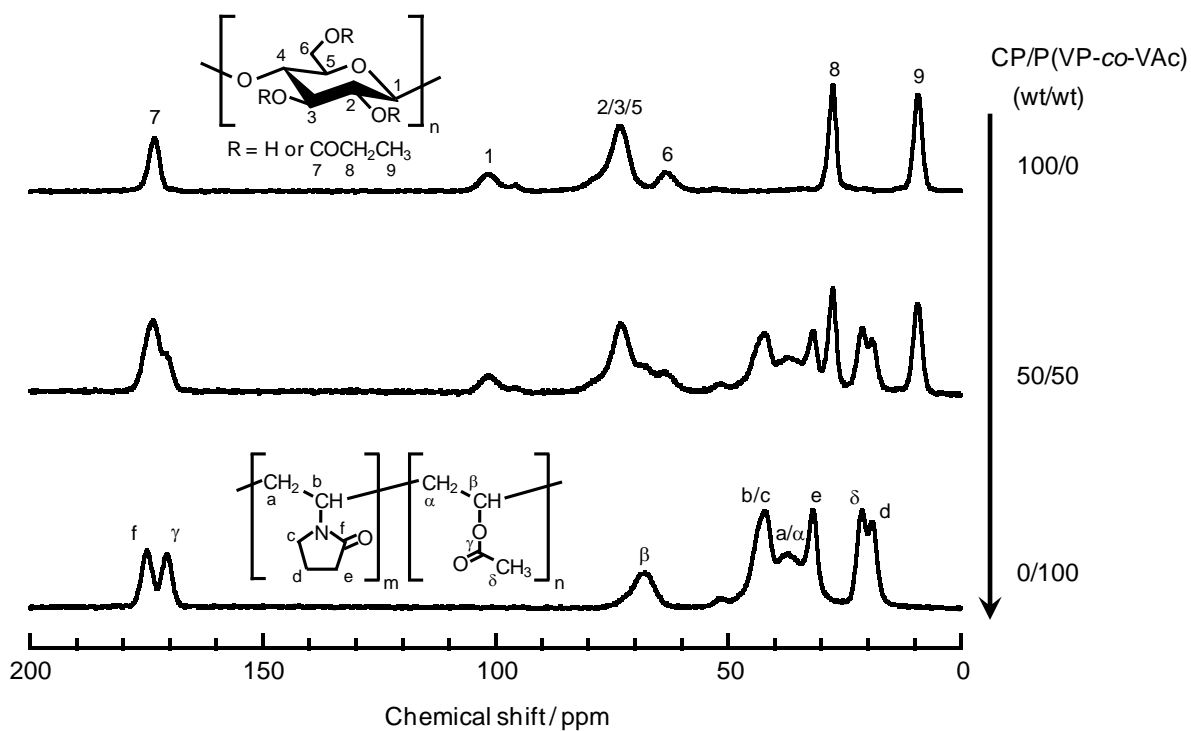
**Fig. 6** DSC thermograms obtained for (a)  $CP_{2.89}/P(VP_{0.52}\text{-}co\text{-}VAc_{0.48})$  and (b)  $CP_{2.89}/P(VP_{0.10}\text{-}co\text{-}VAc_{0.90})$  blends. Arrows indicate a  $T_g$  position.



**Fig. 7** Miscibility map for CP/P(VP-*co*-VAc) blends, as a function of DS of CP and VP fraction in P(VP-*co*-VAc). Symbols indicate that a given pair of CP/P(VP-*co*-VAc) is miscible (○), immiscible (×), or partially miscible (△).

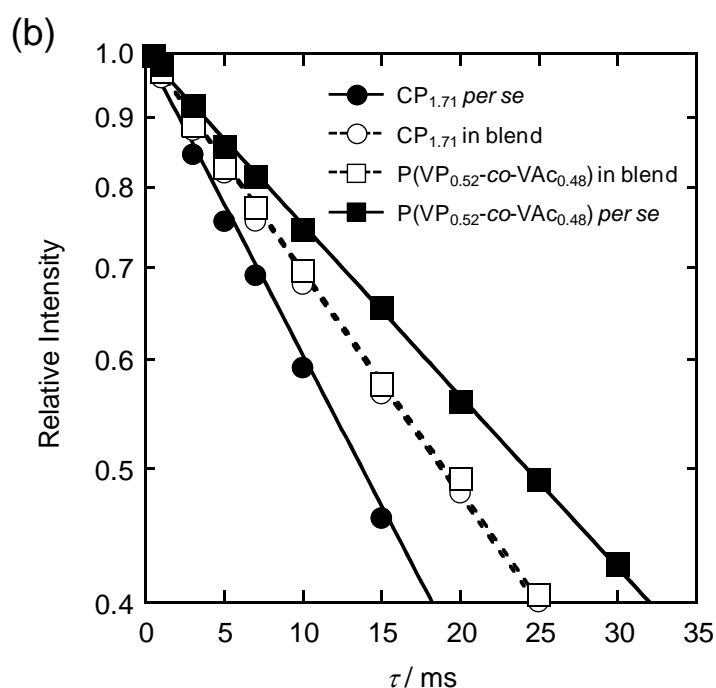
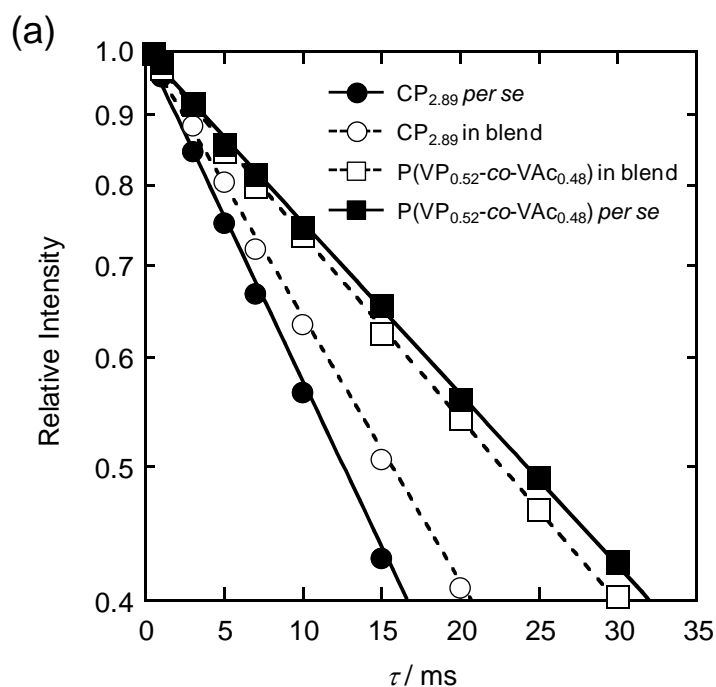


**Fig. 8** Temperature dependence of the dynamic storage modulus  $E'$  and loss modulus  $E''$  for  
(a) CP<sub>2.89</sub>/P(VP<sub>0.10</sub>-co-VAc<sub>0.90</sub>) and (b) CP<sub>2.18</sub>/PVP and CP<sub>2.89</sub>/P(VP<sub>0.52</sub>-co-VAc<sub>0.48</sub>) blends.



**Fig. 9** Solid-state <sup>13</sup>C CP/MAS NMR spectra for CP<sub>2.89</sub>, P(VP<sub>0.52</sub>-co-VAc<sub>0.48</sub>), and their 50/50 blend.





**Fig. 10** Semilogarithmic plots of the decay of  $^{13}\text{C}$  resonance intensities as a function of spin-locking time  $\tau$ , for solid films of (a)  $\text{CP}_{2.89}$ ,  $\text{P}(\text{VP}_{0.52}\text{-co-VAc}_{0.48})$ , and their 50/50 blend, and (b)  $\text{CP}_{1.71}$ ,  $\text{P}(\text{VP}_{0.52}\text{-co-VAc}_{0.48})$ , and their 50/50 blend. The monitoring was conducted for the peak intensity of C2/C3/C5 pyranose carbons of CP and that of  $\text{C}_b/\text{C}_c$  carbons of the copolymer (see Fig. 9).